# Your Guide to Understanding Genetic Conditions

# ANOS1 gene

anosmin 1

### **Normal Function**

The *ANOS1* gene, also known as *KAL1*, provides instructions for making a protein called anosmin-1. This protein is involved in development before birth. Anosmin-1 is found in the extracellular matrix, which is the intricate lattice of proteins and other molecules that forms in the spaces between cells. Anosmin-1 is active in many parts of the developing embryo, including the respiratory tract, kidneys, digestive system, and certain regions of the brain.

Researchers are working to determine the functions of anosmin-1. They have discovered that, in the developing brain, this protein is involved in the movement (migration) of nerve cells and the outgrowth of axons, which are specialized extensions of nerve cells that transmit nerve impulses. The protein also plays a role in regulating contact between nerve cells (cell adhesion).

Anosmin-1 appears to help control the growth and migration of a group of nerve cells that are specialized to process the sense of smell (olfactory neurons). These nerve cells originate in the developing nose and then migrate together to a structure in the front of the brain called the olfactory bulb, which is critical for the perception of odors. Studies suggest that anosmin-1 is also involved in the migration of neurons that produce a hormone called gonadotropin-releasing hormone (GnRH). Like olfactory neurons, GnRH-producing neurons migrate from the developing nose to the front of the brain. GnRH controls the production of several hormones that direct sexual development before birth and during puberty. These hormones are important for the normal function of the ovaries in women and testes in men.

# **Health Conditions Related to Genetic Changes**

# Kallmann syndrome

More than 140 mutations in the *ANOS1* gene have been identified in people with Kallmann syndrome, a disorder characterized by the combination of hypogonadotropic hypogonadism (a condition affecting the production of hormones that direct sexual development) and an impaired sense of smell. This condition can also affect other body systems, and its features vary among affected individuals. Researchers estimate that mutations in the *ANOS1* gene account for 5 to 10 percent of all cases of Kallmann syndrome.

The ANOS1 gene mutations that cause Kallmann syndrome delete part or all of the gene, change single protein building blocks (amino acids) in anosmin-1, or alter the

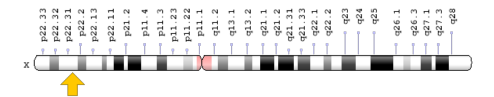
size of the protein. All of these mutations disrupt the normal production or function of anosmin-1 during embryonic development. Researchers suspect that the missing or altered protein is unable to direct the migration of olfactory nerve cells and GnRH-producing nerve cells to their usual locations in the developing brain. If olfactory nerve cells do not extend to the olfactory bulb, a person's sense of smell will be impaired. Misplacement of GnRH-producing neurons prevents the production of sex hormones, which interferes with normal sexual development and causes puberty to be delayed or absent.

It is unclear how *ANOS1* gene mutations lead to other possible signs and symptoms of Kallmann syndrome, including a failure of one kidney to develop (unilateral renal agenesis), hearing loss, and mirror movements of the hands (bimanual synkinesia). Because these features vary among individuals, researchers suspect that other genetic and environmental factors may be involved. Some affected individuals have mutations in one of several other genes in addition to *ANOS1*, and these genetic changes may contribute to the varied features of the condition.

#### **Chromosomal Location**

Cytogenetic Location: Xp22.31, which is the short (p) arm of the X chromosome at position 22.31

Molecular Location: base pairs 8,528,874 to 8,732,187 on the X chromosome (Homo sapiens Annotation Release 108, GRCh38.p7) (NCBI)



Credit: Genome Decoration Page/NCBI

#### Other Names for This Gene

- adhesion molecule-like X-linked
- ADMLX
- anosmin-1
- HHA
- KAL
- KAL1

- KALIG-1
- Kallmann syndrome 1 protein
- Kallmann syndrome protein
- KALM HUMAN
- KMS
- WFDC19

# **Additional Information & Resources**

#### **Educational Resources**

- Endocrinology: An Integrated Approach (first edition, 2001): GnRH and the control
  of gonadotrophin synthesis and secretion
  https://www.ncbi.nlm.nih.gov/books/NBK29/#A1061
- Neuroscience (second edition, 2001): The Olfactory Bulb https://www.ncbi.nlm.nih.gov/books/NBK11158/

## GeneReviews

 Isolated Gonadotropin-Releasing Hormone (GnRH) Deficiency https://www.ncbi.nlm.nih.gov/books/NBK1334

## Scientific Articles on PubMed

PubMed

https://www.ncbi.nlm.nih.gov/pubmed?term=%28%28KAL1%5BTIAB%5D%29+OR+%28ANOS1%5BTIAB%5D%29+OR+%28anosmin+1%5BTIAB%5D%29%29+AND+english%5Bla%5D+AND+human%5Bmh%5D+AND+%22last+1800+days%22%5Bdp%5D

#### **OMIM**

 KAL1 GENE http://omim.org/entry/300836

# Research Resources

- Atlas of Genetics and Cytogenetics in Oncology and Haematology http://atlasgeneticsoncology.org/Genes/GC\_ANOS1.html
- ClinVar https://www.ncbi.nlm.nih.gov/clinvar?term=ANOS1%5Bgene%5D
- HGNC Gene Family: Fibronectin type III domain containing http://www.genenames.org/cgi-bin/genefamilies/set/555

- HGNC Gene Family: WAP four-disulfide core domain containing http://www.genenames.org/cgi-bin/genefamilies/set/361
- HGNC Gene Symbol Report http://www.genenames.org/cgi-bin/gene\_symbol\_report?q=data/ hgnc\_data.php&hgnc\_id=6211
- NCBI Gene https://www.ncbi.nlm.nih.gov/gene/3730
- UniProt http://www.uniprot.org/uniprot/P23352

# **Sources for This Summary**

- Cariboni A, Pimpinelli F, Colamarino S, Zaninetti R, Piccolella M, Rumio C, Piva F, Rugarli EI, Maggi R. The product of X-linked Kallmann's syndrome gene (KAL1) affects the migratory activity of gonadotropin-releasing hormone (GnRH)-producing neurons. Hum Mol Genet. 2004 Nov 15;13(22): 2781-91. Epub 2004 Oct 7.
  - Citation on PubMed: https://www.ncbi.nlm.nih.gov/pubmed/15471890
- Choy C, Kim SH. Biological actions and interactions of anosmin-1. Front Horm Res. 2010;39:78-93. doi: 10.1159/000312695. Epub 2010 Apr 8. Review.
   Citation on PubMed: https://www.ncbi.nlm.nih.gov/pubmed/20389087
- GeneReview: Isolated Gonadotropin-Releasing Hormone (GnRH) Deficiency https://www.ncbi.nlm.nih.gov/books/NBK1334
- Hardelin JP, Julliard AK, Moniot B, Soussi-Yanicostas N, Verney C, Schwanzel-Fukuda M, Ayer-Le Lievre C, Petit C. Anosmin-1 is a regionally restricted component of basement membranes and interstitial matrices during organogenesis: implications for the developmental anomalies of X chromosome-linked Kallmann syndrome. Dev Dyn. 1999 May;215(1):26-44.
   Citation on PubMed: https://www.ncbi.nlm.nih.gov/pubmed/10340754
- Hardelin JP, Levilliers J, del Castillo I, Cohen-Salmon M, Legouis R, Blanchard S, Compain S, Bouloux P, Kirk J, Moraine C, et al. X chromosome-linked Kallmann syndrome: stop mutations validate the candidate gene. Proc Natl Acad Sci U S A. 1992 Sep 1;89(17):8190-4.
   Citation on PubMed: https://www.ncbi.nlm.nih.gov/pubmed/1518845
   Free article on PubMed Central: https://www.ncbi.nlm.nih.gov/pmc/articles/PMC49883/
- Legouis R, Hardelin JP, Levilliers J, Claverie JM, Compain S, Wunderle V, Millasseau P, Le Paslier D, Cohen D, Caterina D, et al. The candidate gene for the X-linked Kallmann syndrome encodes a protein related to adhesion molecules. Cell. 1991 Oct 18;67(2):423-35.
   Citation on PubMed: https://www.ncbi.nlm.nih.gov/pubmed/1913827
- de Castro F, Esteban PF, Bribián A, Murcia-Belmonte V, García-González D, Clemente D. The adhesion molecule anosmin-1 in neurology: Kallmann syndrome and beyond. Adv Neurobiol. 2014; 8:273-92. Review.
  - Citation on PubMed: https://www.ncbi.nlm.nih.gov/pubmed/25300141

Reprinted from Genetics Home Reference: https://ghr.nlm.nih.gov/gene/ANOS1 Reviewed: December 2016 Published: March 21, 2017

Lister Hill National Center for Biomedical Communications U.S. National Library of Medicine National Institutes of Health Department of Health & Human Services